

CLAIMS AMENDMENT

1. (currently amended): A method for producing a soluble protein domain comprising:
(a) ~~preparing two or more DNA fragments by partially digesting a DNA coding for a protein;~~

~~[(b)]~~ (a) expressing at least two nucleotide sequences each encoding a fusion protein which is coded on each of said DNA fragments fused with a DNA encoding a functional comprised of a fragment of a starting protein and a protein exhibiting a function,

~~[(e)]~~ (b) selecting ~~[[the]]~~ a fusion protein exhibiting said function from among two or more fusion the proteins synthesized in step (b) step (a), as comprising a fragment of said starting protein that is a soluble domain, and

~~[(d)]~~ (c) synthesizing the soluble protein domain included in the fusion protein selected in step (b) in a cell-free system, wherein said soluble protein domain is included in said fusion protein selected in step (c).

2. (canceled)

3. (currently amended): The method of claim 1, wherein said ~~functional~~ protein exhibiting a function in step (b) step (a) is any one selected from the group consisting of an enzyme, a binding protein, a luminescent protein and a fluorescent protein, ~~[(or a)]~~ and functional portions thereof.

4. (currently amended): The method of claim 3, wherein said fluorescent protein is a green fluorescent protein or a ~~derivative~~ variant thereof.

5. (currently amended): The method of claim 1, wherein said ~~selection~~ selecting in step (c) step (b) is performed by transforming a recipient in cells with each of containing said DNA fragments and the DNA of said functional protein, nucleotide sequences and by selecting ~~[(the)]~~ a clone which exhibits said function in the obtained transformants.

6. (currently amended): The method of claim 5, wherein said ~~recipient~~ cells are *Escherichia coli* (*E. coli*).

7. (currently amended): The method of claim 1, wherein the fusion proteins are ~~synthesized~~ expressed in a cell-free system, and wherein said ~~selection~~ selecting in step (e) step (b) is performed by measuring the function of the ~~expressed~~ fusion proteins.

8-9. (canceled)

10. (currently amended): A method for producing a soluble protein domain comprising:

(a) ~~constructing a~~ providing an expression vector which expresses a fusion protein of a first protein with second protein that is a green fluorescent protein or a derivative variant thereof, ~~wherein said expression vector comprises a DNA coding for a protein and a gene for said green fluorescent protein or a derivative thereof,~~

(b) ~~preparing two or more DNA fragments by partially digesting said expression vector with DNA decomposing enzyme to obtain two or more DNA fragments of said vector~~ containing deletions of the nucleotide sequence encoding the first protein,

(c) transforming ~~*Escherichia coli*~~ *E. coli* with each of said DNA fragments prepared in step (b) to obtain two or more transformed *E. coli*,

(d) isolating a transformed clone that emits fluorescence among the transformed *E. coli* thus identifying a clone containing DNA that encodes a fusion protein with a soluble protein domain,

(e) recovering the DNA from the isolated transformed clone, and

(f) synthesizing the soluble protein domain ~~which is coded~~ encoded on the recovered DNA in a cell-free system.

11. (currently amended): A method for producing a soluble protein domain comprising:

(a) selecting a fusion protein that exhibits a function characteristic of a functional protein from a plurality of fusion proteins ~~containing~~ each composed of a first protein which is

said functional protein exhibiting a function and a second protein which is a candidate soluble domain, wherein in the selected protein said second protein is a soluble domain, and

(b) synthesizing a soluble ~~protein~~ domain ~~from that was included in the fusion protein~~ selected from step (a).

12. (currently amended): The method of ~~claim 10~~ claim 11, wherein said ~~functional protein contains a second protein is~~ encoded by a DNA fragment ~~of~~ resulting from a partially digested DNA.

13. (currently amended): A method for producing a soluble protein domain comprising:

(a) ~~preparing~~ providing an expression vector comprising a DNA ~~coding for~~ encoding a fusion protein comprised of a first protein and a DNA coding for a second protein which is functional ~~protein~~;

(b) treating said vector ~~by using with~~ a decomposing enzyme and forming to form two or more digested vectors, each vector comprising a fragment of said DNA ~~coding for a protein and the DNA coding for a functional~~ encoding the second protein;

(c) expressing ~~{{a}}~~ fusion protein which is coded on each of said vectors fused with a DNA encoding a functional protein exhibiting a function proteins encoded on the digested vectors obtained in step (b);

(d) selecting the fusion protein exhibiting ~~{{said}}~~ the function characterizing the functional protein among two or more fusion proteins synthesized in step (c) as comprising a soluble domain of said first protein; and

(e) synthesizing the soluble ~~protein~~ domain included in the fusion protein selected in step (d) in a cell-free system, ~~wherein said soluble protein domain is included in said fusion protein selected in step (d)~~.

14. (currently amended): The method of claim 13, wherein the ~~selection~~ selecting of step (d) is performed by transforming ~~a recipient cell~~ cells with the ~~expression vector comprising each of said DNA fragments and the DNA of said functional protein~~ digested vectors, and selecting ~~the a~~ a clone which exhibits said function in the obtained transformants.

15. (new): A method to synthesize a soluble domain that is a portion of a starting protein which method comprises synthesizing, in a cell-free system, a protein identified as said soluble domain by:

- (a) preparing a multiplicity of fusion proteins, each said fusion protein comprising a functional portion and a fragment of said starting protein,
- (b) assessing each fusion protein for the function of the functional portion; and
- (c) identifying, as a soluble domain, fragments of said protein which are contained in fusion proteins that exhibit the function of the functional portion.

16. (new): The method of claim 15, wherein said preparing is performed in a cell-free system.

17. (new): The method of claim 15, wherein said preparing is performed intracellularly.

18. (new): The method of claim 17, wherein said preparing is performed *in vivo* in *E. coli*.

19. (new): The method of claim 15, wherein the functional portion comprises an enzyme, a binding protein, a luminescent protein or a fluorescent protein or functional portions thereof.

20. (new): The method of claim 19, wherein the fluorescent protein is green fluorescent protein or a variant thereof.

21. (new): A method to produce a soluble domain that is a portion of a starting protein which method comprises

(a) expressing, in each of at least two *E. coli* colonies, a fusion protein comprising green fluorescent protein (GFP) or a variant thereof fused to a fragment of said starting protein and

(b) identifying a transformed *E. coli* colony that emits fluorescence, whereby a colony comprising a fusion protein containing a fragment that is a soluble domain is identified, and

(c) producing the soluble protein domain identified in step (b).

22. (new): The method of claim 21, wherein each said fragment is obtained by a process comprising digesting nucleic acid encoding a fusion protein comprising said GFP or variant and said starting protein with a DNA digesting enzyme.

23. (new): The method of claim 22, wherein said digesting is in only either from the 3' or 5' end of the nucleic acid.